Effervescent tablets are an interesting pharmaceutical/nutraceutical dosage form that offers some unique advantages when compared with simple tablets. However, the manufacturing process involves some critical steps that need to be addressed carefully during formulation and factory design.
**Fundamentals of Effervescents**

Oral dosage forms are still the most popular way of taking medication, despite having some disadvantages compared with other methods. One such disadvantage is the risk of slow absorption of the active pharmaceutical ingredient (API), which can be overcome by administering the drug in liquid form and, therefore, possibly allowing the use of a lower dosage. Liquid dosage forms also benefit from higher levels of patient compliance, especially pleasant-tasting ones, and overcome any potential issues with dysphagia in geriatric and pediatric populations. However, because many APIs only show a limited level of stability in liquid form, effervescent tablets, which are dissolved in water before administration, have been formulated as an alternative dosage form.

Advantages of effervescent tablets compared with other oral dosage forms include

- an opportunity for formulators to improve the taste
- a gentler action on a patient’s stomach
- marketing aspects (fizzy tablets may have more consumer appeal than traditional dosage forms).

The disadvantages of effervescent dosage forms are the need for larger tablets, a complex production process and, very often, the need for specialist packaging materials.

Effervescents generally comprise at least one organic acid and an alkali metal carbonate salt. In certain formulations, a second organic acid is sometimes added. Carbon dioxide is formed if this mixture comes into contact with water. Typical examples of the acids and alkalis used include the following (also see Figure 1):

- citric acid
- tartaric acid
- malic acid
- fumaric acid
- adipic acid
- sodium bicarbonate/carbonate/sesquicarbonate
- potassium bicarbonate/carbonate.

**Production**

Producing effervescent tablets requires a conventional solid dosage form manufacturing process that has been adapted to include additional features because of the unique characteristics of the product.

**Material handling:** The primary material used in the manufacture of effervescents is relatively hygroscopic; that is, it absorbs moisture from the air. However, this must be prevented because it will initiate the effervescent reaction. One of the principle strategies used to overcome this problem is a completely closed material handling system, which includes intermediate bulk containers (IBCs), docking stations and split valve technology. In addition, the ventilating air must contain a sufficiently low moisture content. This method is particularly useful if potent actives are handled that require a high level of operator protection.

**Granulation and drying:** Because most tablets are compressed by high-speed rotary tablet presses, the material fed into the presses has to have properties that prevent segregation and ensure homogenous filling of the dies — to produce tablets of equal weight and API content. The most common approach to achieving materials with these characteristics is to granulate the raw materials. Because wet granulation will initiate the effervescent reaction, several alternatives have been established.¹

**Dry methods:** Dry methods, such as slugging, direct compression and roller compaction are regularly used to produce solid dosage forms. These are the preferred methods of producing effervescents because no liquid is involved, which means that no additional drying step is required. Another advantage is the reduced need for equipment because of the limited number of unit operations required and, as a consequence, ventilation of the machinery and/or the facility can be simplified. Indeed, the roller compaction method, if properly automated, can lead to very high throughput. However, the main arguments against the use of dry methods are the need for expensive excipients, which is especially critical as effervescent tablets tend to be large. Another disadvantage is that by using methods such as roller compaction, the flowability of the material (into the dies of the tablet press) can be improved ... but not the compressibility.

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![Figure 1](image.png)

1 mol R-COOH + 1 mol NaHCO₃ → 1 mol R-COONa + 1 mol H₂O or

1 mol R-COONa + 2 mol H₂O + 1 mol CO₂ → 1 mol R-COONa + 1 mol H₂O + 1 mol CO₂
**Wet granulation:** For the wet granulation process, two separate granulation steps are run (one each for the alkaline and the acid components) with a subsequent dry blending step. This can be done in a high shear granulator (with subsequent drying), a single pot or in a fluid bed spray granulator. The advantage of this method is that only conventional equipment is needed, which could also be used for granulating and drying other materials.

The major disadvantages are the running time required and cleaning, particularly if two parallel lines are not available for the two granulations. The blending process can be a critical step and may affect the homogeneity of the tablets because not all the materials are bonded into one granule as in a conventional wet granulation process.

**Organic solvents:** Because the effervescent reaction is only started if the materials come into contact with water — and not if they come into contact with organic solvents — one possibility is to use such solvents as a granulation fluid. This can be done in a high shear granulator (with subsequent drying), a single pot or in a fluid bed spray granulator.

The method offers a number of advantages, which originate from the lower heat of evaporation when compared with water, high throughput, the possibility of drying at lower temperatures and the freedom to use a number of different excipients to achieve the desired product characteristics.

The only disadvantage of this method is the need for more complex equipment to handle the fluids. If a fluid bed is used, then a complex system for exhaust gas treatment is required because the mixture of organic vapor and a large amount of non-condensable process gas has to be treated. This would not be the case in a tray dryer or a single pot as only the organic vapor must be handled. Also, additional care is required to address potential safety issues, such as operator protection and explosion prevention.

**Water:** It is possible to use water as a granulation fluid. Only a very small amount of water is added, which initiates a pre-effervescent reaction. This results in some carbon dioxide being produced, as well as water, which consequently acts as the granulation fluid, producing more carbon dioxide and more water. The cycle must be stopped at a certain point by starting the drying process and removing the water more quickly than it’s produced by the effervescent reaction.

This can be done using a high shear granulator with subsequent drying by discharging the material into a pre-heated fluid bed dryer at the end of the granulation process. The most critical steps are discharge and transfer; hence, this method works well for small and medium batch sizes. However, it may lead to problems for larger batch sizes because of the long time needed for this operation.

A second variation is to use a single pot, in which the granulation process can be curtailed by switching to the drying mode. Drying, in this situation, can be enhanced by using a double jacket and vacuum system or by using gas or microwave technology. Again, this would be acceptable for small or medium batches but, because of a poor surface volume ratio, would be too slow for larger batches. Only the use of microwaves allows the use of this technology at larger scales (1200 L).

**Fluid bed spray granulation:** Fluid bed spray granulation is a unique process in which granulation and drying take place simultaneously. This ensures a constant low moisture level, limiting the pre-effervescent reaction to a minimum. In addition, when using a fluid bed for drying, it is easy to achieve a very low final moisture level for storage. The disadvantage is that more granulation fluid is needed than in a high shear process.

Recent developments in continuous processing allow the use of additional methods. Dry raw materials can be fed into a twin-screw granulator. Here, because the residence time of the material in the granulator is less than one minute, a water-based binder fluid can be used. Subsequently, the material is immediately transferred to a subsequent dryer.

**Lubricants**

It is common practice in tablet production to add a lubricant after granulation; the most commonly used substance is magnesium stearate. Its function is to improve the flow of the material, which is extremely important because the dies of a tablet press are filled by volume. A second function is to prevent the tablet sticking to the punch faces or to the walls of the dies.

During effervescent production, substances such as magnesium stearate should not be used because they are insoluble in water and, consequently, a film will form on top of the water after the tablet has dissolved. Strategies to overcome this problem are the use of other lubricants that are soluble in water (such as a mixture of spray dried L-leucine and polyethylene glycol).
Tablet Compression

The compression of effervescent tablets is different from the compression of normal tablets. For long-term storage, a very low moisture content is required — typically less than 0.3% water compared with approximately 2% for traditional tablets. An additional drawback is that, typically, effervescent formulations only contain a small amount of binder to facilitate rapid dissolution in water.

Sometimes the post-granulation drying step is stopped at a moisture content of 2–3% to safeguard trouble-free compression. To ensure ongoing stability, the tablets then need to be dried to a moisture content of 0.3%. Additionally, it’s important that there is a seamless connection between the drying step, the compression step and the post-drying step as any delays could destabilize the formulation.

Additionally, effervescent tablets are usually quite large, which often leads to insufficient tablet hardness and, consequently, broken or damaged tablets. This, in turn, results in a poor yield and a need to stop the press or packaging line. One way of overcoming this problem is to increase the dwell time by modifying the pre-compression assembly of the tablet press.

If working without a lubricant, the poorer flow characteristics of the material can be addressed using a constant level powder feed system. This consists of a rotary valve that guarantees a constant powder pressure on the forced filling station, which, in connection with two independently driven feed wheels, will ensure accurate die filling.

The second problem, when working without a lubricant, is that the tablets may stick to the die walls or on the punch faces. This issue can be overcome using a punch face and die wall lubrication system, which allows the addition of a very small amount of solid or liquid lubricant to the punch faces and the die walls just before they come in contact with the granules. It should be noted that because of their design, tablet presses easily accommodate the processing of effervescent materials and only release dry air into the compression zone, removing the need to vent the entire room.

Packaging

After the material has been pressed into tablets, the surface area of the material has been significantly reduced, which means that the rate at which moisture is absorbed from the air has also been reduced. Consequently, the dehumidification of the environmental air is now less critical.

Blisters and tube arrangements are used for packaging; for example, standard packaging materials are used in the packaging of food products or some nutraceuticals when shelf-life is not critical. This is not acceptable for pharmaceutical products. Aluminum, which has low water permeability, is used instead of standard polymer blister materials. If ten or more individual tablets are packed into one tube, very dry air can be added; but, as the patient opens the tube to take out the first tablet, ambient moist air will enter and destroy the remaining effervescent tablets. To overcome this, silica gel or other drying agents are incorporated into most tube lids.

Conclusion

Effervescents are an interesting pharmaceutical dosage form offering some unique advantages compared with traditional tablets. The manufacturing process involves some critical steps that need to be addressed carefully during formulation and factory design.

References

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